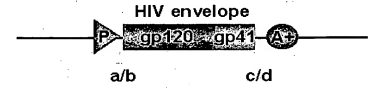
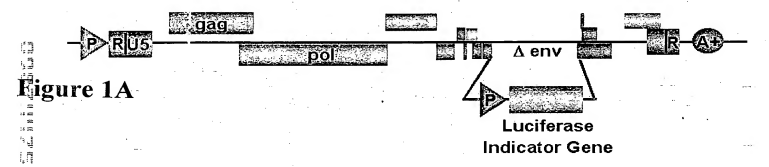
METHODS

Envelope Expression Vector: pHIVenv

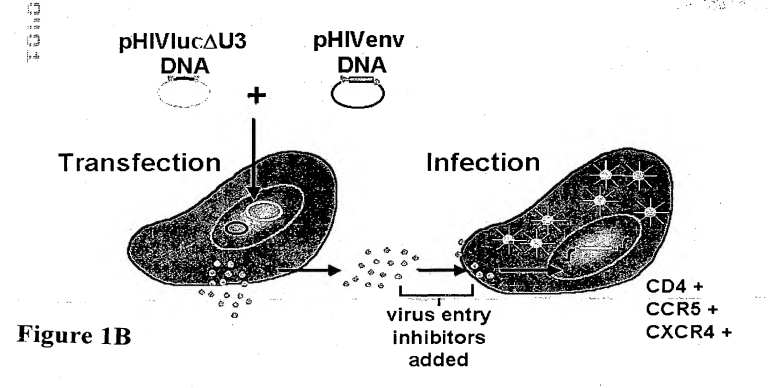
1. .



HIV-1 Expression Vector: pHIVIucΔU3

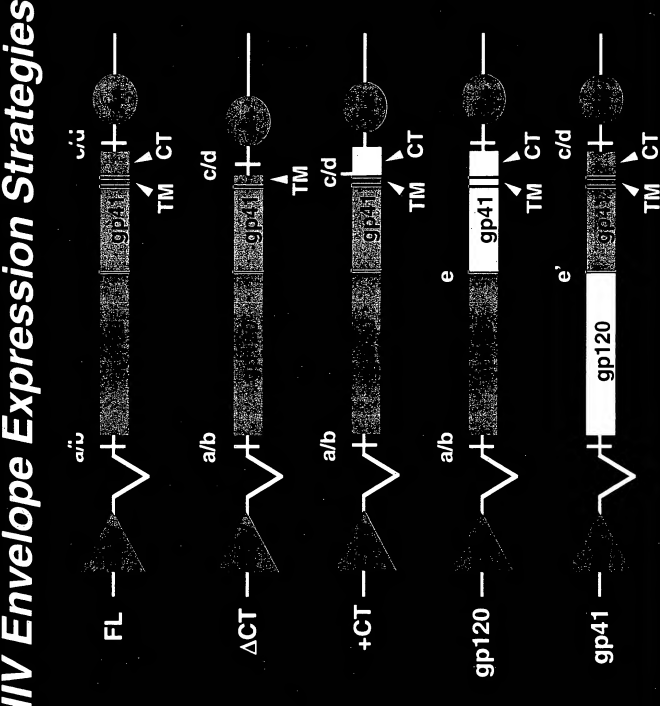


PhenoSense HIV Entry Cell Assay



1

HIV Envelope Expression Strategies



Co-Receptor Tropism Screen

CCR5-expressing cells

STATE OF THE

No drug inhibitor **CCR5** CXCR4

- TOTAL 的数

3963

345 Sept

inhibitor

Replicate 2 Replicate 1

No drug

がある。 | Control | Cont 2.7 CANADA RAMAS the second second

CXCR4-expressing cells

7

55 AN 1245

CXCR4

inhibitor

inhibitor

CCR5

N. N. S.

<100 RLU

100-1000 RLU

1000-10,000 RLU >10,000 RLU

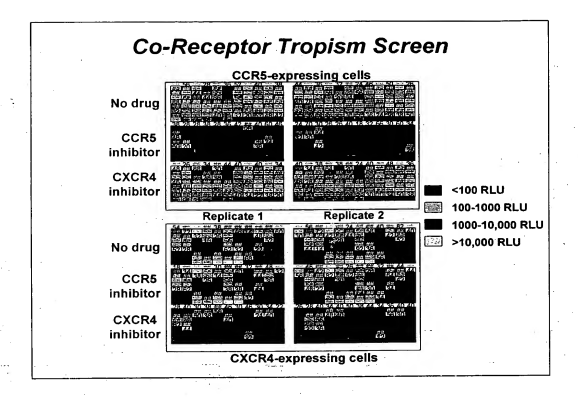


Figure 3A. Co-receptor Tropism Screening Assay

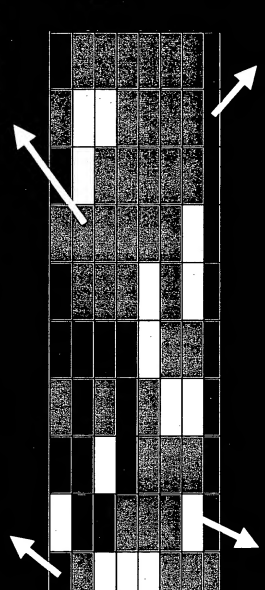
In this embodiment, the assay is performed using two cell lines. One cell line expresses CD4 and CCR5 (top six panels). The other cell line expresses CD4 and CXCR4 (bottom six panels). The assay is performed by infecting cells with a large number of recombinant virus stocks derived from cells transfected with pHIVenv and pHIVlucΔU3 vectors. Tthe example shown represents the analysis of 96 viruses formatted in a 96 well plate. Infections are performed in the absence of drug (no drug), or in the presence of a drug that preferentially inhibits either R5 tropic (CCR inhibitor) or X4 tropic (CXCR4 inhibitor) viruses. Co-receptor tropism is assessed by comparing the amount of luciferase activity produced in each cell type, both in the presence and absence of drug (see Figure 3B for interpretation of assay results).

Co-Receptor Tropism Assay Interpretation

R5:X4 R5 cells X4 cells no drug R5 inhibitor X4 inhibitor %inhib by R5 inhibitor %inhib by X4 inhibitor

R5 X4 R5:X4
no drug CASSI CASSI CASSI
R5 inhibitor CASSI CASSI CASSI
%inhib by R5 inhibitor CASSI CASS

21



active RLU limit: 100 tropism ratio limit: 5

DUAL or MIXED

NON-VIABLE

R5 X4 R5:X4
no drug 43 42
R5 inhibitor
X4 inhibitor
X4 inhibitor
%inhib by R5 inhibitor

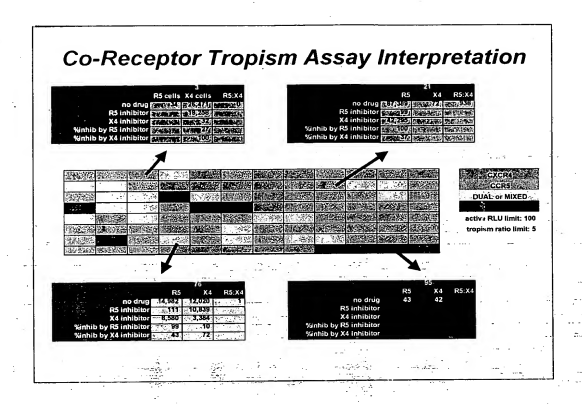


Figure 3B

Determining co-receptor tropism.

In this embodiment, the results of the assay are interpreted by comparing the ability of each sample virus to infect (produce luciferase activity) in cells expressing CD4/CCR5 (R5 cells) or cells expressing CD4/CXCR4 (X4 cells). The ability of a CCR5 or CXCR4 inhibitor to specifically block infection (inhibit luciferase activity) is also evaluated.

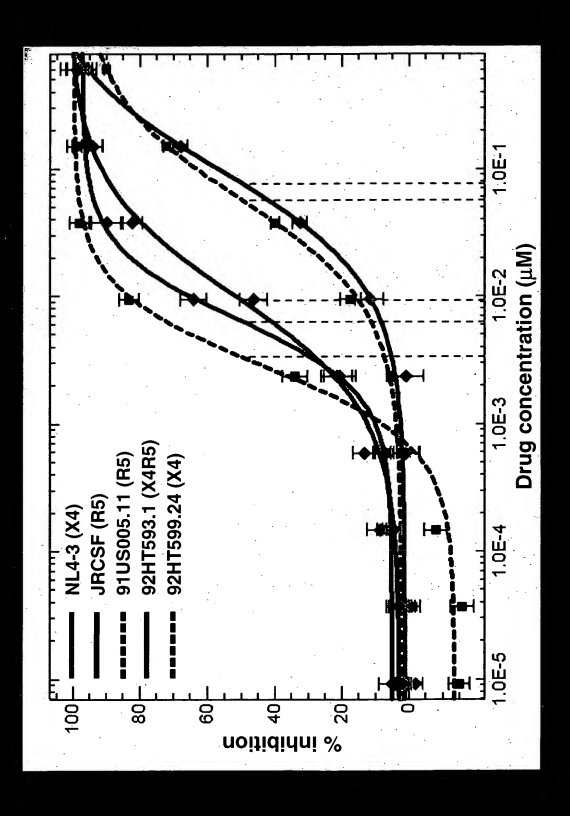
X4 tropic viruses (green panels)- infect X4 cells but not R5 cells. Infection of X4 cells is blocked by the CXCR4 inhibitor.

R5 tropic viruses (blue panels)- infect R5 cells but not X4 cells. Infection of R5 cells is blocked by the CCR5 inhibitor.

Dual tropic or X4/R5 mixtures (yellow panels)- infect X4 and R5 cells. Infection of R5 cells is blocked by the CCR5 inhibitor and infection of X4 cells is blocked by the CXCR4 inhibitor.

Non-viable viruses (red panels)- do not replicate in either X4 or R5 cells.

Entry Inhibitor Susceptibility: Fusion Inhibitor



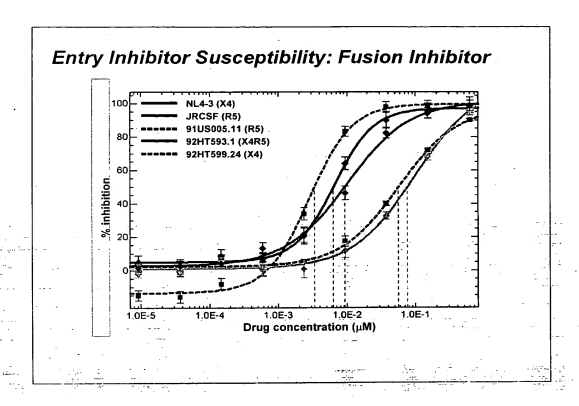


Figure 4A.

Measuring Entry Inhibitor Susceptibility: Fusion Inhibitor

In this embodiment, susceptibility to the fusion inhibitor T-20 is demonstrated. Cells expressing CD4, CCR5 and CXCR4 were infected in the absence of T-20 and over a wide range of T-20 concentrations (x-axis log10 scale). The percent inhibition of viral replication (y-axis) was determined by comparing the amount of luciferase produced in infected cells in the presence of T-20 to the amount of luciferase produced in the absence of T-20. R5 tropic, X4 tropic and dual tropic viruses were tested. Drug susceptibility is quantified by determining the concentration of T-20 required to inhibit 50% of viral replication (IC50, shown as vertical dashed lines). Viruses with lower IC50 values are more susceptible to T-20 than viruses with higher IC50 values.

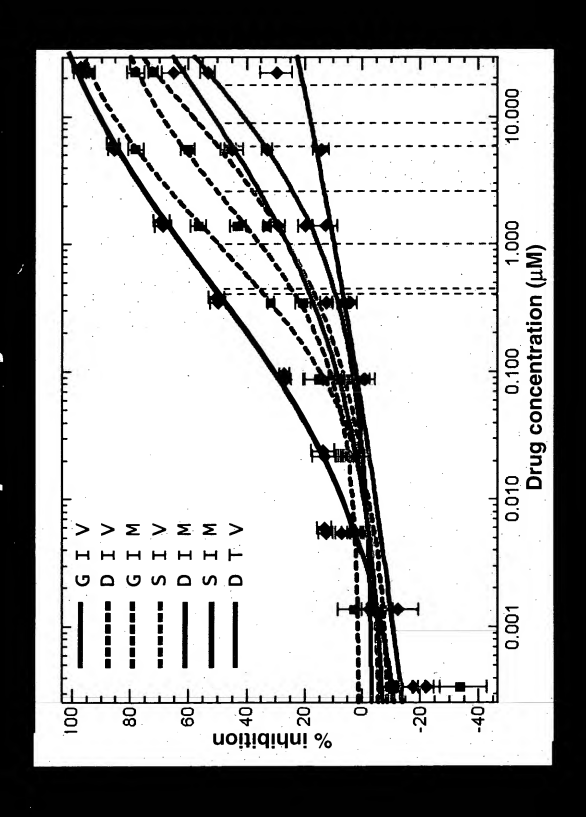
NL4-3: well-characterized X4 tropic strain

JRCSF: well-characterized R5 tropic strain

91US005.11: R5 tropic isolate obtained from the NIH AIDS Research and Reference Reagent Program (ARRRP)

92HT593.1: Dual tropic (X4R5) isolate obtained from the NIH ARRRP.

Reduced Susceptibility: Fusion Inhibitor



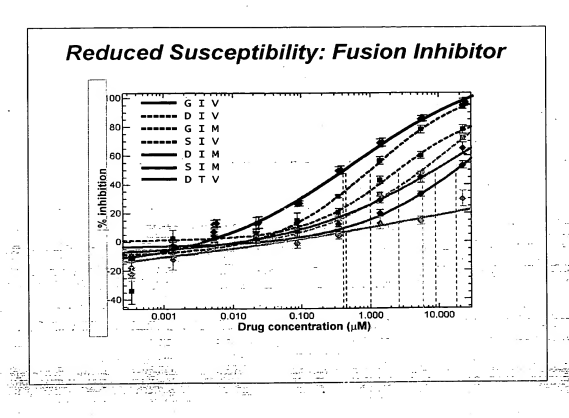


Figure 4B.

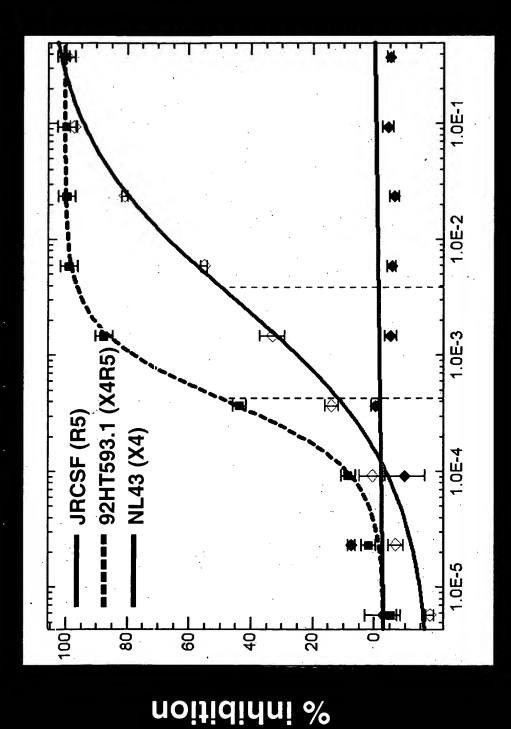
Measuring Entry Inhibitor Susceptibility: Drug Resistance Mutations

In this embodiment, reduced susceptibility to the fusion inhibitor T-20 conferred by specific drug resistance mutations in the gp41 envelope protein is demonstrated. Cells expressing CD4, CCR5 and CXCR4 were infected in the absence of T-20 and over a wide range of T-20 concentrations (x-axis log10 scale). The percent inhibition of viral replication (y-axis) was determined by comparing the amount of luciferase produced in infected cells in the presence of T-20 to the amount of luciferase produced in the absence of T-20. Isogenic viruses containing one or two specific mutations in the gp41 transmembrane envelope protein were tested (highlighted in red in the figure legend). Drug susceptibility is quantified by determining the concentration of T-20 required to inhibit 50% of viral replication (IC50, shown as vertical dashed lines). Viruses with lower IC50 values are more susceptible to T-20 than-viruses with higher IC50 values.

No mutation (wildtype sequence): GIV

Single mutations: GIV, DIM, SIV

Double mutations: DIM, SIM, DTV



Drug: R5 Inhibitor Cell: CD4/CCR5

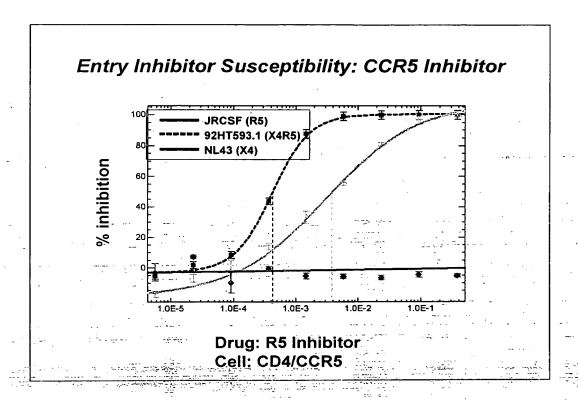


Figure 5A.

Measuring Entry Inhibitor Susceptibility: CCR5 Inhibitor

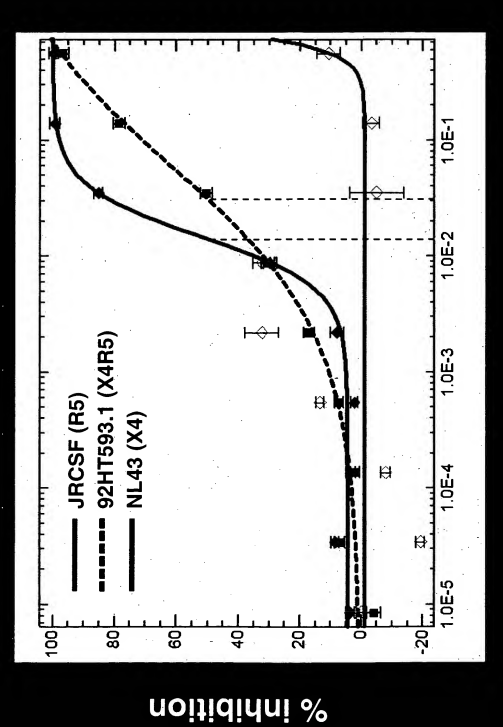
In this embodiment, susceptibility to a CCR5 inhibitor (merck compound) is demonstrated. Cells expressing CD4 and CCR5 (R5 cells) were infected in the absence of the CCR5 inhibitor and over a wide range of CCR5 inhibitor concentrations (x-axis log10 scale). The percent inhibition of viral replication (y-axis) was determined by comparing the amount of luciferase produced in infected cells in the presence of CCR5 inhibitor to the amount of luciferase produced in the absence of CCR5 inhibitor. R5 tropic, X4 tropic and dual tropic viruses were tested. Drug susceptibility is quantified by determining the concentration of CCR5 inhibitor required to inhibit 50% of viral replication (IC50, shown as vertical dashed lines). Viruses with lower IC50 values are more susceptible to the CCR5 inhibitor than viruses with higher IC50 values. The X4 tropic virus did not infect the R5 cells.

NL4-3: well-characterized X4 tropic strain

JRCSF: well-characterized R5 tropic strain

92HT593.1: Dual tropic (X4R5) isolate obtained from the NIH ARRRP.

Entry Inhibitor Susceptibility: CXCR4 Inhibitor



Drug: X4 Inhibitor Cell: CD4/CXCR4

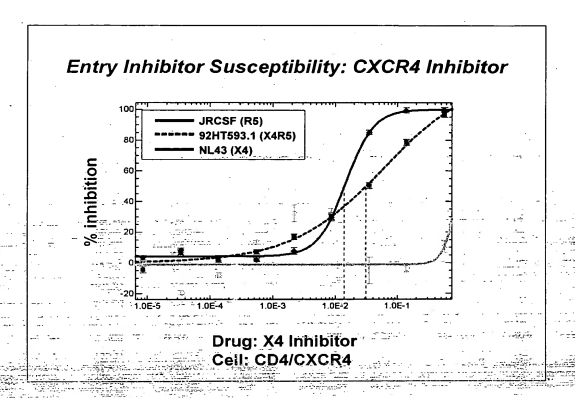


Figure 5B.

Measuring Entry Inhibitor Susceptibility: CXCR4 Inhibitor

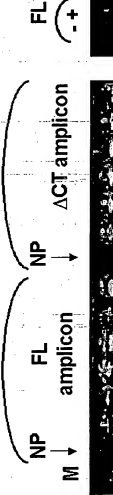
In this embodiment, susceptibility to a CXCR4 inhibitor (AMD3100) is demonstrated. Cells expressing CD4 and CXCR4 (X4 cells) were infected in the absence of the CXCR4 inhibitor and over a wide range of CXCR4 inhibitor concentrations (x-axis log10 scale). The percent inhibition of viral replication (y-axis) was determined by comparing the amount of luciferase produced in infected cells in the presence of CXCR4 inhibitor to the amount of luciferase produced in the absence of CXCR4 inhibitor. R5 tropic, X4 tropic and dual tropic viruses were tested. Drug susceptibility is quantified by determining the concentration of CXCR4 inhibitor required to inhibit 50% of viral replication (IC50, shown as vertical dashed lines). Viruses with lower IC50 values are more susceptible to the CCR5 inhibitor than viruses with higher IC50 values. The R5 tropic virus did not infect the X4 cells.

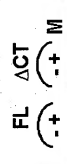
NL4-3: well-characterized X4 tropic strain

JRCSF: well-characterized R5 tropic strain

92HT593.1: Dual tropic (X4R5) isolate obtained from the NIH ARRRP.

Envelope Sequence Amplification





1. R5

3. R5/ 3. R5/ 5. R5/ 5. R5

6. R5/X4 NP: HIV negative plasma

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16 July 16	高峰			# of isolates	15	3
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	ر کلکات کامسوکات			5 6	5 6 # of isolates	6 # of isolates

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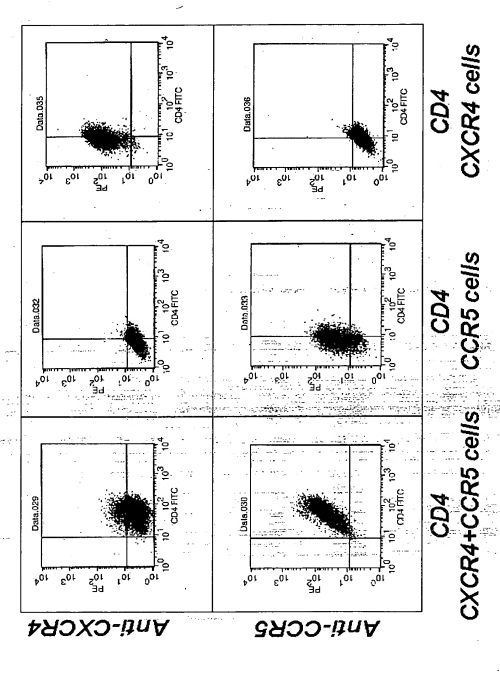
3

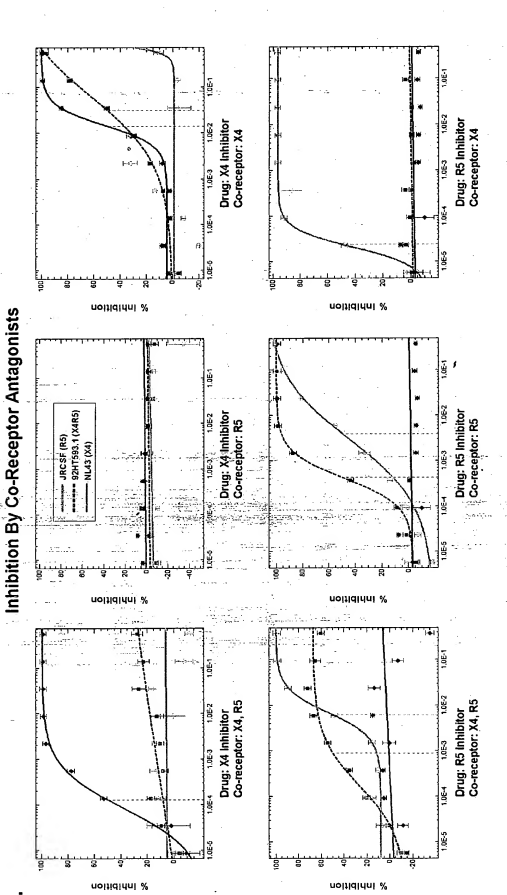
Co-Receptor Tropism	# of isolates
 X	15
R5	24
X4/R5	15
 Undefined	35
Envelope Subtype	# of isolate
Clade A	2
Clade B	92
Clade C	_
Clade D	_
Clade E	ෆ
-	

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Target Cell Receptor and Co-Receptor Expression





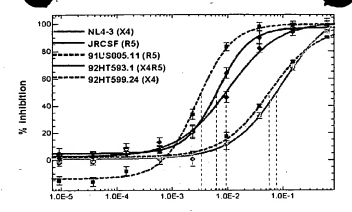
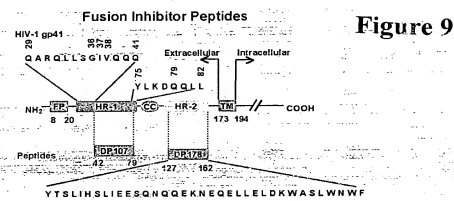
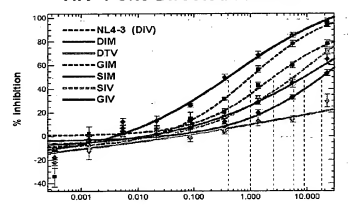


Figure 4A



Rimsky, et al., J. Virol. 72 (2):986-993

HIV-1 Site Directed Mutants



I IZUIC TD	Fig	ure	4B
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SDM Virus	DP 178 Sens.a	Fold Changeb
HXB2 GIV	DI 170 SCHS.	1.0
NL4-3 G I V	S	5.2
NL4-3 D I V	5	12.8
NL4-3 S I V	S	74.2
NL4-3 G I M	S	33.0
NL4-3 D I M	R	113.0
NL4-3 S I M	R	227.4
NL4-3 D T V	R	>281.8
JRCSF G I V		2.1
JRCSF D I V		104.0

^a Rimsky et al., J. Virol. 72(2):986-993 ^b Fold change in IC50 (vs. HXB2) using PhenoSense HIV Entry Assay